=> d his

(FILE 'CAPLUS' ENTERED AT 12:27:37 ON 27 FEB 2004)
DELETE HIS

FILE 'REGISTRY' ENTERED AT 13:09:55 ON 27 FEB 2004
L1 STRUCTURE UPLOADED
L2 1 S L1
L3 14 S L1 SSS FULL
L4 13 S L3 NOT C32 H26 N2 O2/MF
L5 12 S L4 NOT C34 H31 N3 O4/MF
L6 11 S L5 NOT C35 H30 N2 O4 /MF

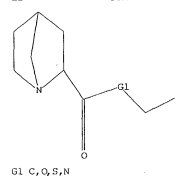
FILE 'CAPLUS' ENTERED AT 13:13:17 ON 27 FEB 2004 L7 4 S L6

FILE 'REGISTRY' ENTERED AT 13:44:38 ON 27 FEB 2004

FILE 'MARPAT' ENTERED AT 13:44:50 ON 27 FEB 2004

L8 0 S L3
L9 21 S L3 SSS FULL
L10 16 S L9/COMPLETE
L11 16 S L10 NOT L7
L12 0 S L11 AND NEURONAL

=> d 11 L1 HAS NO ANSWERS



Structure attributes must be viewed using STN Express query preparation.

=> d 111 1-16 bib abs

ANSWER 1 OF 16 MARPAT COPYRIGHT 2004 ACS on STN 140:5203 MARPAT ΑN Preparation of opioid and opioid-like compounds for therapeutic uses TI TN Yen, Mao-Hsiung; Fan, Chin-Tsai PA. Jenken Bioscience, Inc., USA SO PCT Int. Appl., 58 pp. CODEN: PIXXD2 DТ Patient. LA English FAN.CNT 1 PATENT NO. KIND DATE APPLICATION NO. DATE WO 2003097608 20031127 WO 2003-US15461 20030516 PΙ A2 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC,

GW, ML, MR, NE, SN, TD, TG

NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ,

PRAI US 2002-380841P 20020517

(+)-Morphinan derivs., such as I [R = H, allyl, Ph, benzyl, alkyl, heteroarylalkyl, heterocyclyl, etc.], were prepared for therapeutic use in pharmaceutical compns. These morphinans are claimed for use in the treatment of neurodegenerative diseases, such as Parkinson's disease, Alzheimer's disease, cognition deficit, memory loss, and stroke, cancers, such as skin cancer, small cell lung cancer, testicular cancer, esophageal cancer, breast cancer, endometrial cancer, ovarian cancer, central nervous system cancer, liver cancer, and prostate cancer, cardiac disorders, such as cardiac ischemia, congestive heart failure, and hypertension, as well as for the treatment of septic shock, inflammation, organ damage and neurol. disorders. Thus, (+)-3-methoxy-17-allylmorphinan hydrobromide was prepared in 63% yield via a an allylation reaction (+)-3-methoxymorphinan hydrochloride with allyl bromide using Et3N in THF followed by treatment of the base I (R = allyl) thus formed with HBr. Pharmacol. activities were determined by measurement of TNF- α , IL-10, nitrate, serum glutamic oxaloacetic transaminase, serum glutamic pyruvic transaminase and creatinine levels.

L11 ANSWER 2 OF 16 MARPAT COPYRIGHT 2004 ACS on STN

AN 136:86056 MARPAT

ΤI Preparation of amino acid-N-carboxy anhydrides having substituent at nitrogen

IN Tsunoda, Hidetoshi; Sekiguchi, Michiru; Iizuka, Hajime; Sakai, Kazuya

PA Mitsui Chemicals, Inc., Japan

PCT Int. Appl., 91 pp. SO

CODEN: PIXXD2

DT Patent

LA Japanese

FAN.	CNT	1																
	PATENT NO.			KII	ND	DATE			A	PPLI	CATI	ON NO	o.	DATE				
PI	WO 2002002538			A1 20020110				WO 2001~JP5780			0	20010704						
		W:	BR,	CA,	CN,	IN,	KR,	US										
		RW:	AT,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,
			PT,	SE,	TR													
	JP 2002201183		A2 20020716			J!	20	01-2	0397	1	2001	0704						
	EP 1298127			A1 20030402			EP 2001-949905			5	20010704							
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
			ΙE,	FI,	CY,	TR												
	US 2002173664		A.	11 20021121			US	3 20	02-7	0156		2002	0304					
	US 6670447		·B	2	2003	1230												
PRAI	JP 2000-201745		20	0007	04													

JP 2000-303522 20001003

WO 2001-JP5780 20010704

OS CASREACT 136:86056

GI

EP 750908

Al

```
Amino acid-N-carboxy anhydrides having an N-acyl substituent at nitrogen
      which are represented by the following general formula [I; R1 = CH3,
      (CH3)2CH, (CH3)3C, C6H5CH2, C6H5CH2OCOCH2CH2; R2 = C6H5, CH3, CH3(CH2)8, C6H5CH2CH2] and a process for producing the same. These compds. easily
      react with free amino acids, alcs. or nucleophilic agents such as anions
      and serve as useful intermediates in producing amino acid derivs.,
      optically active compds., peptides, polypeptides and the like, which are useful in various fields including medicines and pesticides, at a high
      yield. Moreover, a process for producing diamides R2CONHCHR1CONR3R4 (R1,
      R2 as above; R3 = H, alkyl, cycloalkyl, aryl) by using title compds. I
      with amine derivs. represented by R3NHR4. These diamides are also
      appropriately usable in producing amino acid derivs., optically active
      compds., peptides, polypeptides and the like.
THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 7
                  ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 3 OF 16 MARPAT COPYRIGHT 2004 ACS on STN
      134:95517 MARPAT
AN
      Quinuclidine derivatives for the treatment of neurological disorders
TТ
      Lauffer, David; Mullican, Michael
IN
      Vertex Pharmaceuticals Incorporated, USA
٩٩
      PCT Int. Appl., 55 pp.
SO
      CODEN: PIXXD2
DT
      Patent
      English
LA
FAN.CNT 1
                                                       APPLICATION NO. DATE
      PATENT NO.
                            KIND DATE
                            A1 20010111
                                                       WO 2000-US18355 20000705
      WO 2001002405
ΡI
           W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BB, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YUL, ZA, ZW, AM, AZ, RY, KG, KZ, MD, RIL, TJ, TM
           XU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

1200440

A1 20020502

EP 2000-945145 20000705
       EP 1200440
           R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL 2003503500 T2 20030128 JP 2001-507841 20000705
       JP 2003503500
                                    20020905
                                                       US 2002-39886
                                                                              20020103
       US 2002123507
                              A1
                              B2
                                    20031209
      US 6660748
                                                        US 2003-632618
                                                                              20030801
                                    20040212
      US 2004029912
                             A1
PRAI US 1999-142509P 19990706
       WO 2000-US18355 20000705
                             20020103
      Quinuclidine derivs. are provided for treating or preventing neuronal
       damage associated with neurol. diseases. The invention also provides compns.
       comprising the compds. of the invention and methods of using those compns.
       for treating or preventing neuronal damage.
                  THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 10
                  ALL CITATIONS AVAILABLE IN THE RE FORMAT
      ANSWER 4 OF 16 MARPAT COPYRIGHT 2004 ACS on STN
       124:15483 MARPAT
AN
       Remedy for infectious diseases
ΤI
       Suzuki, Fujio
       Tsumura and Co., Japan
 PΑ
       PCT Int. Appl., 53 pp.
SO
       CODEN: PIXXD2
 DΤ
       Patent
       Japanese
 LA
 FAN.CNT 1
                             KIND DATE
                                                        APPLICATION NO. DATE
       PATENT NO.
                             ----
                                                        WO 1995-JP491
                                                                              19950317
                              A1
                                    19950928
       WO 9525517
 PΙ
            W: AU, CA, CN, JP, KR, US
            RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
                                                        CA 1995-2185825 19950317
       CA 2185825
                              AA 19950928
                                                                              19950317
                                   19951009
                                                        AU 1995-19605
       AU 9519605
                              A1
       AII 689235
                              B2
                                     19980326
                                   19970102
                                                         EP 1995-912470 19950317
```

```
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE
                                         CN 1995-192984 19950317
                           19970416
    CN 1147766
                      Α
                           19990601
    US 5908857
                      Α
                                          US 1996-704673
                                                            19960918
                                          US 1998-135591
                                                            19980818
    US 6030980
                      Α
                           20000229
PRAI JP 1994-72820
                     19940318
    JP 1994-72821
                     19940318
    WO 1995-JP491
                     19950317
    An infectious disease remedy comprises at least one member selected from
    the group consisting of Aconitum root alkaloids (e.g. benzoylmesaconine
    and 14-anisoylaconine), Aconitum roots and exts. thereof, gingerols and
    analogs thereof, and ginger rhizomes and products of treatment thereof.
    It has a remarkable effect of restoring the protective activity against
    infections and is useful for treating and preventing viral infection,
     fungal infection, and opportunistic infection. Benzoylmesaconine
     (10µq/kq/day) administered orally to cytomegalovirus-infected mice with
    exptl. burn effectively controlled the infection. Tablets were formulated
    containing corn starch 198.5, light anhydrous silicic acid 1, and
    14-anisoylaconine 0.5 g.
    ANSWER 5 OF 16 MARPAT COPYRIGHT 2004 ACS on STN
T.1.1
ΑN
    123:339729 MARPAT
    Preparation of N-(carbamoylcyclohexyl)indole-3-carboxamides and analogs as
    tachykinin antagonists
    Sisto, Alessandro; Fincham, Christopher; Potier, Edoardo; Manzini,
IN
    Stefano; Arcamone, Federico; Lombardi, Paolo
    A. Menarini Industrie Farmaceutiche Riunite S.R.L., Italy; Malesci
     Istituto Farmacobiologico S.P.A.
SO
    PCT Int. Appl., 36 pp.
    CODEN: PIXXD2
DT
     Patent
LA
    English
FAN.CNT 1
                      KIND DATE
                                           APPLICATION NO. DATE
     PATENT NO.
                            _____
                                                           19941202
                                           WO 1994-EP4012
РΤ
     WO 9515311
                      A1
                           19950608
        W: AM, AU, BB, BG, BR, BY, CA, CN, CZ, EE, FI, GE, HU, JP, KE, KG,
            KP, KR, KZ, LK, LR, LT, LV, MD, MG, MN, MW, NO, NZ, PL, RO, RU,
             SD, SI, SK, TJ, TT, UA, US, UZ, VN
         RW: KE, MW, SD, SZ, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU,
            MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN,
             TD, TG
                            19950608
                                           CA 1994-2177994 19941202
     CA 2177994
                       AA
                                           AU 1995-12731
                                                            19941202
                            19950619
     AU 9512731
                       A1
                                           EP 1995-903789
                                                            19941202
                       A1
                            19960918
     EP 731790
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE
                                           JP 1994-515404
                                                            19941202
                      T2
                            19970624
     JP 09506348
                                           US 1996-656068
                                                            19960531
                            19980602
     US 5760248
                       А
                      19931203
PRAI IT 1993-FI247
     WO 1994-EP4012
                      19941202
GT
          R2
```

AB Title compds. [I; A,B = N, CH; R1,R2 = H, OH, halo, Me(CH2CH2O)2, etc.; R3,R4,R8 = H, alkyl; R3R4 = (CH2)1-3; R5,Y = (hetero)aryl(alkyl), etc.; X = CO2R6, CH2OR6, NR7COR6, etc.; R6,R7 = groups cited for R5; Z = CH2, CO; dashed line = optional bond] were prepared Thus, aminocyclohexanecarboxylic acid II (R = OH) was amidated by (S)-H2NCH(CH2Ph)CO2CH2Ph to give II [R = (S)-NHCH(CH2Ph)CO2CH2Ph]. II [R = (R)-NHCHR5NMeCOCHMePh; R5 = 2-naphthylmethyl] had pKi of 8.7 for antagonism of substance P in vitro.

L11 ANSWER 6 OF 16 MARPAT COPYRIGHT 2004 ACS on STN

AN 122:284619 MARPAT

GΙ

```
Preparation of substituted triazole and tetrazole derivatives as
TI
     insecticides.
     Dick, Michael R.
     DowElanco, USA
PA
     U.S., 10 pp.
SO
     CODEN: USXXAM
     Patent
DT
     English
LA
FAN.CNT 1
                                             APPLICATION NO. DATE
                       KIND DATE
     PATENT NO.
                       ____
                                                               19930826
                                             US 1993-112498
     US 5393767
                        А
                             19950228
РΤ
                                             WO 1994-US10703 19940921
                             19960328
                        A1
     WO 9608968
         W: AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, ES, FI, GB, HU,
              JP, KR, KZ, LK, LU, LV, MG, MN, MW, NL, NO, NZ, PL, PT, RO, RU,
             SD, SE, SI, SK, UA, UZ
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
                             19960409
                                             AU 1994-78003
                                                               19940921
     AU 9478003
                       A1
PRAI US 1993-112498
                       19930826
     WO 1994-US10703 19940921
GΙ
     The title compds. I and II (Y = CH, N; Z = H, F, Cl, Br, CN, CONH2,
AB
     alkoxycarbonyl, etc;R = N-containing heterocyclyl) are prepared as insecticides
     and acaricides, especially active against sucking insects, such as brown plant
     hoppers, and phytophagous mites, such as two-spotted spider mites.
     Exo-3-(5-aminotetrazol-2-yl)-1-azabicyclo[2.2.1]heptane is typical.
     ANSWER 7 OF 16 MARPAT COPYRIGHT 2004 ACS on STN
Lll
     122:133852 MARPAT
AN
      Preparation of peptide analog tachykinin antagonists.
TΙ
     Arcamone, Federico; Lombardi, Paolo; Manzini, Stefano; Potier, Edoardo;
IN
      Sisto, Alessandro
     A. Menarini Industrie Farmaceutiche Riunite S.R.L., Italy; Malesci
      Istituto Farmacobiologico S.P.A.
      PCT Int. Appl., 39 pp.
 SO
      CODEN: PIXXD2
 DΤ
      Patent
      English
 LA
 FAN.CNT 1
                                              APPLICATION NO.
                                                                DATE.
      PATENT NO.
                       KIND DATE
      _____
                       ____
                                              WO 1993-EP3387
                                                                19931202
                             19940623
      WO 9413694
                        A1
          W: AU, BB, BG, BR, CA, CZ, FI, HU, JP, KP, KR, KZ, LK, LV, MG, MN,
              MW, NO, NZ, PL, RO, RU, SD, SK, UA, US, UZ, VN
          RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
                                              CA 1993-2151062 19931202
                        AA 19940623
      CA 2151062
                              19940704
                                              AU 1994-56508
                                                                19931202
      AU 9456508
                         A1
                                              EP 1994-901948
                                                                19931202
                              19950920
      EP 672052
                         Α1
                              19970716
      EP 672052
                         В1
          R: DE, ES, FR, GB
                         T3 19971016
                                              ES 1994-901948
                                                                19931202
      ES 2105605
                                              US 1995-448460
                                                                19950602
                              19970624
      US 5641802
                         Α
 PRAI IT 1992-MI2779
                        19921204
      WO 1993-EP3387
                        19931202
```

Title compds. [I; Y = H, (substituted) alkyl, alkenyl, alkylnyl, cycloalkyl, etc.; R1, R2 = H, OH, halo; R1R2 = O; A, B = N, CH; R3, R4 = H, alkyl, alkenyl, alkynyl; R3R4 = (CH2)n; n = 1-3; R5 = alkyl, aryl, alkylaryl, arylalkyl; R6, R7 = H, alkyl, aryl, arylalkyl, alkylaryl; dotted line = optional double bond], were prepared Thus, Me trans-2-aminocyclohexanecarboxylate (preparation given) was coupled with indolin-3-carboxylic acid using hydroxybenzotriazole, 1-ethyl-3-(3'-dimethylaminopropyl)carbodiimide, and diisopropylethylamine in CH2Cl2 to give Me N-(indolin-3-ylcarbonyl)-trans-2-aminocyclohexanecarboxylate. This was saponified with aqueous NaOH and the product was condensed with phenylalanine N-methyl-N-benzylamide (preparation given) using bromotripyrrolidinephosphonium hexafluorophosphate and diisopropylethylamine in CH2Cl2 to give the N-methyl-N-benzylamide of Na-[[N-(indolin-3-ylcarbonyl)(R,R)-trans-2-amino]cyclohexanoyl]phenylalanine and the corresponding diastereomer. I inhibited Substance P binding to guinea pig ileum by 95-100% at 1 μ M.

```
ANSWER 8 OF 16 MARPAT COPYRIGHT 2004 ACS on STN
T.1.1
     122:106537 MARPAT
ΑN
     Preparation of tetrazolyl peptide analogs as fibrinogen receptor
ΤI
     antagonists
     Nutt, Ruth F.; Veber, Daniel F.
IN
     Merck and Co., Inc., USA
PA
so
     PCT Int. Appl., 53 pp.
     CODEN: PIXXD2
DT
     Patent
     English
LA
FAN.CNT 1
                                               APPLICATION NO. DATE
     PATENT NO.
                     / KIND DATE
                               _____
                                                _____
                              19940428
                                               WO 1993-US9569 19931005
     WO 9409029
                         A1
PΤ
         W: AU, BB, BG, BR, BY, CA, CZ, FI, HU, JP, KR, KZ, LK, LV, MG, MN,
              MW, NO, NZ, PL, RO, RU, SD, SK, UA, US
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG 19940823 US 1992-961221 19921014
     HS 5340798
                                               AU 1994-53229
                                                                   19931005
                               19940509
     AU 9453229
                         A1
                        19921014
PRAI US 1992-961221
     WO 1993-US9569
                        19931005
GT
```

AB AECH2CONHCHBCONHCHDD1 (A = primary, secondary, or tertiary amino group; B = carboxy or thiol group; D = H, alkyl, heteroalkyl, aryl, heteroaryl; Dl = 5-substituted tetrazolyl; E = nitrogen, carbocyclyl, heterocyclyl, carboaryl, heteroaryl group), were prepared as inhibitors of integrin protein complex function relating to cell attachment activity. Thus, title compound (I) (solution phase preparation given) inhibited platelet aggregation with IC50 = 0.011 nM.

WO 1994-JP218

GΙ

```
ANSWER 9 OF 16 MARPAT COPYRIGHT 2004 ACS on STN
     121:280563 MARPAT
     Preparation of quinuclidine derivatives as cerebral function ameliorants
ΤI
     Fuse, Yoshihide; Yamamoto, Kozo; Kishida, Hideyuki; Miwa, Toshiaki;
TN
     Hidaka, Takayoshi; Katsumi, Ikuo
     Kanegafuchi Kagaku Kogyo Kabushiki Kaisha, Japan
PΑ
     PCT Int. Appl., 31 pp.
SO
     CODEN: PIXXD2
DT
     Patent
LA
     Japanese
FAN.CNT 1
     PATENT NO.
                                           APPLICATION NO.
                                                             DATE
                      KIND DATE
                            _____
     _____
                                                             19940214
     WO 9419348
                       A1
                            19940901
                                           WO 1994-JP218
PI
        W: US
        RW: CH, DE, FR, GB, IT
                                                             19930217
                      A2 19940830
                                           JP 1993-28281
     JP 06239861
                                           EP 1994-906386
                                                            19940214
     EP 638569
                       A1
                          19950215
        R: CH, DE, FR, GB, IT, LI
5494917 A 19960227
                                           US 1994-313118
                                                            19941012
     US 5494917
PRAI JP 1993-28281
                      19930217
```

19940214

The title compds. I [X = (CH2)a, etc.; a = 0 - 4; Y = (CH2)b, etc.; b = 0 - 3; Z = (CH2)c, etc.; c = 0 - 3; n = 1 - 5; R1 = H, (un)substituted Ph, etc.; R2 = H, alkyl, etc.] are prepared Quinuclidine derivs. II (cis) and III (trans) were prepared from 3-quinuclidinone and di-Et m-phenoxybenzylphosphonate. In in vitro tests for affinity for M1 and M2 receptors using (3H)-pirenzepine and (3H)-N-methylscopolamine, resp., III showed IC50 of 23±3 nM (M1 receptors) and IC50 of 4400±100 nM (M2 receptors). Formulations containing I are given.

DATE

19920722

```
L11 ANSWER 10 OF 16 MARPAT COPYRIGHT 2004 ACS on STN
     121:83070 MARPAT
AN
    Azabicyclic tachykinin receptor antagonists
TΙ
    Swain, Christopher John
IN
    Merck Sharp and Dohme Ltd., UK
PΑ
    Brit. UK Pat. Appl., 25 pp.
SO
     CODEN: BAXXDU
DT
     Patent
LA
     English
FAN.CNT 1
     PATENT NO.
                                           APPLICATION NO.
                      KIND DATE
                                           GB 1992-15527
     GB 2268931
                      A1
                            19940126
```

19920722

$$\bigcap_{N=1}^{Q} \bigvee_{Y=1}^{X-R^2}$$

GT

PRAI GB 1992-15527

```
Title compds. [I; O = residue of an azabicyclic ring system; R1 =
     (halo)phenyl, trifluoromethylphenyl; R2 = (substituted)phenyl; X = 0, S,
     CH2, etc.; Y = H, OH, halo, etc.] are claimed as tachykinin receptor
     antagonists (no data). No prepared I are reported.
    ANSWER 11 OF 16 MARPAT COPYRIGHT 2004 ACS on STN
L11
     119:243594 MARPAT
AN.
     Preparation of substituted oxadiazole and thiadiazole compounds as
TΙ
     acaricides and insecticides.
     Dick, Michael R.; Chang, Chi Ping; Dripps, James E.; Wollowitz, Susan
     DowElanco, USA
PA
     PCT Int. Appl., 29 pp.
SO
     CODEN: PIXXD2
DT
     Patent
     English
T.A
FAN.CNT 1
                                             APPLICATION NO. DATE
     PATENT NO.
                       KIND
                            DATE
                                             _____
                       ____
                             _____
                                            WO 1992-US10493 19921204
                             19930805
                       A1
PΤ
     WO 9314636
         W: AT, AU, BB, BG, BR, CA, CH, CS, DE, DK, ES, FI, GB, HU, JP, KR,
         LK, LU, MG, MN, MW, NL, NO, NZ, PL, PT, RO, RU, SD, SE, UA
RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
             BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, SN, TD, TG
                                             US 1992-824658
                                                             19920123
                             19930914
     US 5244906
                       Α
                                             CA 1992-2105556 19921204
                            19930724
     CA 2105556
                       AA
                                             AU 1993-32396
                                                              19921204
     AU 9332396
                        A1
                             19930901
                        B2
                             19940721
     AU 651516
                                                              19921204
                                             EP 1993-900881
     EP 577788
                        A1
                            19940112
         R: DE, ES, FR, GB, IT, NL
                                             BR 1992-5804
                                                               19921204
                             19940517
     BR 9205804
                        Α
                                             JP 1992-513191
                                                               19921204
                        T2
                            19941020
     JP 06509359
     CN 1097545
                        Α
                             19950125
                                             CN 1993-109067
                                                              19930723
PRAI US 1992-824658
                       19920123
```

WO 1992-US10493 19921204

The title compds. I, II and III [Y = O, S; Z = H, F, Cl, Br, CONH2, CO2R1, OR1, SR1, NH2, (un)substituted Me or Et, etc.; R = N-containing heterocyclyl; R1 = Me or Et] are prepared as.acaricides and insecticides. The cyclization of Me 1-azabicyclo[2,2,1]heptane-3-carboxylate with acetamide oxime, in the presence of NaOEt, gave 3-(3-methyl-1,2,4-oxadiazol-5-yl)-1-azabicyclo[2,2,1]heptane (IV). IV gave >50% control of the 3rd instar green leafhopper (Nephotettix cincticeps), at >2 ppm, and of brown planthopper (Nilaparvata lugens) nymphs, at >0.5 ppm, in laboratory expts.

```
L11 ANSWER 12 OF 16 MARPAT COPYRIGHT 2004 ACS on STN AN 119:139139 MARPAT TI Azabicyclic compounds as tachykinin antagonists
```

IN Ladduwahetty, Tamara; Swain, Christopher J.

PA Merck Sharp and Dohme Ltd., UK

SO Eur. Pat. Appl., 21 pp. CODEN: EPXXDW

DT Patent LA English

FAN.CNT 1 DATE APPLICATION NO. PATENT NO. KIND DATE EP 1992-201931 19920627 Al 19930414 EP 536817 PΤ R: CH, DE, FR, GB, IT, LI, NL CA 1992-2072676 19920629 CA 2072676 AA 19930106 US 1992-905974 19920629 19931026 US 5256671 Α JP 1992-178314 19920706 JP 06157524 19940603 A2 19970528 JP 2614687 B2 PRAI GB 1991-14551 19910705 GB 1991-14887 19910710 19920303 GB 1992-4578

GΙ

$$R^3$$
 R^4
 R^5
 R^2
 R^2
 R^2
 R^2
 R^2
 R^2
 R^2
 R^2
 R^3
 R^4
 R^2
 R^4
 R^6
 R^6

The azabicyclic compound I (Q = residue of an optionally substituted azabicyclic ring system; the dotted line represents an optional double bond; $\bar{X} = H$, HO, =O, halo; R1 = H, Ph, thienyl which may be substituted by halo or F3C; R2 = Ph, thienyl, benzyl which may be substituted by halo or F3C; R3, R4, R5, = H, C1-6 alkyl, C2-6 alkynyl, C2-6 alkenyl, halo, cyano, alkoxy, alkylthio amino, etc. were prepared as tachykinin antagonists. Thus, 2-(diphenylmethyl)quinuclidin-3-one was treated with disopropyl cyanomethylphosphonate followed by reduction with DIBAL-H to give E-3-(formylmethylene)-2-benzhydrylquinuclidine, which underwent Grignard reaction with bromoanisole to give the E-3-[(2-methoxyphenyl)-2hydroxyethylidene]-2-benzylhydrylquinuclidine II and its stereoisomer. The tachykinin antagonist 1C90 of II was 50 $\eta M.$

ANSWER 13 OF 16 MARPAT COPYRIGHT 2004 ACS on STN

118:80807 MARPAT AN

Preparation of benzyloxyazabicycloalkanes as tachykinin antagonists ΤI

Baker, Raymond; Swain, Christopher; Seward, Eileen M. Merck Sharp and Dohme Ltd., UK IN

PA

Eur. Pat. Appl., 26 pp.

CODEN: EPXXDW

DТ Patent

LA English

FAN.CNT 1				
PATENT NO.	KIND D	ATE	APPLICATION NO	D. DATE
PI EP 499313	A1 1	9920819	EP 1992-20030	3 19920204
EP 499313	B1 1	9970611		
R: AT, BE,	CH, DE,	DK, ES, FR	, GB, GR, IT, LI,	LU, NL, PT, SE
US 5242930	A 1	9930907	US 1992-83082	2 19920204
AT 154354	E 1	9970615	AT 1992-20030	3 19920204
CA 2060949	AA 1	9920812	CA 1992-20609	49 19920210
JP 05078354	A2 1	9930330	JP 1992-25068	19920212
JP 2500279	B2 1	9960529		
PRAI GB 1991-2809	1991021	.1		
GB 1991-7403	1991040	19		
GB 1991-13892	1991062	:7		
GB 1991-14553	1991070	15		
GT				

$$R^{5}$$
 R^{4}
 R^{1}
 R^{2}
 R^{2}
 R^{2}
 R^{3}
 R^{4}
 R^{2}
 R^{2}
 R^{3}
 R^{4}
 R^{5}
 R^{5

Title compds. [I; Q = azabicyclic ring residue X = O, S; Y = H, OH; R1, R2= (halo- or CF3-substituted) Ph, thienyl; R3-R5 = H, alkyl, alkenyl, alkynyl, halo, cyano, NO2, CF3, Me3Si, OR6, SMe, SOMe, SO2Me, NR6R7, NR6COR7, NR6CO2R7, CO2R6, CONR6R7; R6, R7 = H, alkyl, Ph, CF3], were

prepared Thus, cis-2-diphenylmethyl-1-azabicyclo[2.2.2]octane-3-ol (preparation by reduction of the corresponding ketone given) in dimethoxyethane at 0° was treated with 18-crown-6, KN(SiMe3)2 in PhMe, and 3-O2NC6H4CH2Br in dimethoxyethane. The mixture was stirred 1 h to give 50% coupling product, which was converted to the title compound II. II at 1.0 μM gave > 10% inhibition of substance P Me ester-induced contraction of guinea pig ileum longitudinal muscle.

```
ANSWER 14 OF 16 MARPAT COPYRIGHT 2004 ACS on STN
1.11
```

117:198529 MARPAT ΑN

Ophthalmic pharmaceuticals containing substituted pyridine derivatives for treatment of glaucoma

Lotti, Victor; Showell, Graham A. ΤN

Merck Sharp and Dohme Ltd., UK PΑ

Can. Pat. Appl., 25 pp. CODEN: CPXXEB

DΤ Patent

LA English

FAN.CNT 1 PATENT NO.	KIND DATE	APPLICATION NO.	DATE
PI CA 2058249	AA 19911208	CA 1991-2058249	19911220 19911211
EP 492902 R: CH, DE,	Al 19920701 FR, GB, IT, LI, NL		
JP 05140158 PRAI GB 1990-27824	A2 19930608 19901221	JP 1991-361151	19911220
GB 1991-12307	19910607		ber a managem
AB Pyridine derive	., substituted on the	pyridine nucleus	

azabicyclic ring system with >5 ring atom are used in ophthalmic pharmaceuticals for treatment of glaucoma. Thus, 0.05% 3-[2-(6-ethoxypyridin)yl-1-azabicyclo[2,2,2]octane hydrogen oxalate (I) changed intraocular pressure by 3.0 mmHg. An eye drop contained I 0.5, benzalkonium chloride 0.02%, Na2EDTA 0.05, NaCl 0.8, and water to 100%.

```
ANSWER 15 OF 16 MARPAT COPYRIGHT 2004 ACS on STN
L11
```

113:152263 MARPAT

Preparation of 1-azabicycloalkanes as cholinergics TI

Galliani, Giulio; Barzaghi, Fernando; Bonetti, Carla; Toja, Emilio TN

PA Roussel-UCLAF, Fr.

so Eur. Pat. Appl., 13 pp.

CODEN: EPXXDW

DT Patent

LA	French									
FAN.CNT 1										
	PATENT NO.	KIND DATE	APPLICATION NO.	DATE						
PΙ	EP 366561	A2 , 19900	502 EP 1989-402973	19891027						
	EP 366561	A3 19910	918							
	R: AT, BE,	CH, DE, ES,	FR, GB, GR, IT, LI, LU, NL	•						
	JP 02178280	A2 19900	711 JP 1989-277350	19891026						
	US 5015655	A 19910	514 US 1989-426778	19891026						
	CA 2001686	AA 19900	428 CA 1989-2001686	19891027						
	US 5183893	A 19930	202 US 1991-664120	19910326						
PRAI	IT 1988-22452	19881028								
	US 1989-426778	19891026								
GI										

$$\begin{array}{c|c} \text{CH2}_{\text{N}} & \text{CR1} = \text{NOR2} \\ \hline & \text{N} & \text{II} \\ \hline & \text{CHZ} \\ \end{array}$$

The title compds. [I; R1, R2 = H, alkyl, alkenyl, etc.] were prepared AB Hydrolysis-oxidation of (methoxymethylene)quinuclidine II in CHCl3 with HClO4 gave 3-quinuclidinecarboxaldehyde (III, Z = O), which was condensed with

<code>H2NOMe.HCl</code> to give III (Z = NoMe) (IV). IV at 1.10-3 at 1.10-3 and 1.10-8M showed muscarinic and nicotinic effects on guinea pig ileum. Tablets and gelatin capsules containing I were formulated.

```
L11 ANSWER 16 OF 16 MARPAT COPYRIGHT 2004 ACS on STN
    111:153786 MARPAT
ΑN
    Preparation of nonaromatic azacyclic or azabicyclic ring-containing
ΤI
    1,3-oxazoles or 1,3-thiazoles for the treatment of senile dementia
    Baker, Raymond; Snow, Roger J.; Saunders, John; Showell, Graham A.
TN
    Merck Sharp and Dohme Ltd., UK
PΑ
    Eur. Pat. Appl., 31 pp.
    CODEN: EPXXDW
    Patent
DΨ
    English
FAN.CNT 1
    PATENT NO.
                     KIND DATE
                                          APPLICATION NO. DATE
                                          EP 1988-308126 19880901
                           19890315
    EP 307141
                      A2
                      A3 19890412
    EP 307141
                      В1
                           19930113
     EP 307141
```

R: AT, BE, CH, DE, ES, FR, GB, IT, LI, LU, NL, SE 19880901 AT 1988-308126 19930115 AT 84415 F. ES 1988-308126 19880901 ES 2053748 Т3 19940801 DK 1988-5032 19880909 19890428 DK 8805032 A 19890614 JP 1988-225568 19880910 JP 01151576 A2 US 1993-86389 19930701 19940628 US 5324723 Α 19870910 PRAI GB 1987-21343 GB 1988-1759 19880127 EP 1988-308126 19880901 19880902 US 1988-239892 19900205 US 1990-454492 US 1992-912469 19920713

GI For diagram(s), see printed CA Issue.

The title compds. [I; X = O, S; Rl = nonarom. azacyclic or azabicyclo ring; R2 = H, halo, CF3, OH, alkoxy, cyano, (substituted) amino, hydrazino, etc.] useful for treating mental illness (e.g. senile dementia) and as analgesics, were prepared Saponification of 3-methoxycarbonylquinuclidine with LiOH.H2O in refluxing MeOH and methylation of the resulting 3-carboxyquinuclidine Li salt with MeLi in THF gave 3-acetylquinuclidine which was heated 6 h with iodine and H2NCSNH2 on a steam bath to give 3-(2-amino-1,3-thiazol-4-yl)quinuclidine. Treatment of this in 50% hypophosphorus acid with aqueous NaNO2 for 5 days gave 3-(1,3-thiazol-4-yl)quinuclidine (II). Tablets containing II.hydrogen oxalate 25.0, microcryst. cellulose 37.25, modified food corn starch 37.25, and Mg stearate 0.50 mg were formulated.

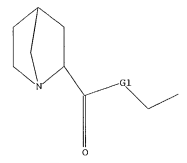
=>

```
10387318
```

=> d his

```
(FILE 'CAPLUS' ENTERED AT 12:27:37 ON 27 FEB 2004)
DELETE HIS
```

```
FILE 'REGISTRY' ENTERED AT 13:09:55 ON 27 FEB 2004
L1
                STRUCTURE UPLOADED
              1 S L1
L2
             14 S L1 SSS FULL
1.3
             13 S L3 NOT C32 H26 N2 O2/MF
L4
             12 S L4 NOT C34 H31 N3 O4/MF
             11 S L5 NOT C35 H30 N2 O4 /MF
L6
     FILE 'CAPLUS' ENTERED AT 13:13:17 ON 27 FEB 2004
L7
              4 S L6
=> d 11
L1 HAS NO ANSWERS
                STR
```



G1 C, O, S, N

Structure attributes must be viewed using STN Express query preparation.

```
=> d 1-4 bib abs hitstr
```

```
ANSWER 1 OF 4 CAPLUS COPYRIGHT 2004 ACS on STN
L7
      2004:41475 CAPLUS
ΑN
DN
      140:111404
      Preparation of N-aryl diazaspirocyclic compounds as nicotinic cholinergic
TΤ
      receptor modulators for treating nervous system and other disorders
      Bhatti, Balwinder S.; Miller, Craig H.; Schmidt, Jeffrey D.
      Targacept, Inc., USA
PA
      PCT Int. Appl., 101 pp.
SO
      CODEN: PIXXD2
DΤ
      Patent
      English
LA
FAN.CNT 1
                                                       APPLICATION NO. DATE
      PATENT NO.
                            KIND DATE
                                    20040115
                                                       WO 2003-US20524 20030627
PΙ
      WO 2004005293
                             A2
           W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
                CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
                LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
                PL, PT, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
           RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ,
                 GW, ML, MR, NE, SN, TD, TG
PRAI US 2002-394337P
                                    20020705
GΙ
```

RN

CN

Compds., pharmaceutical compns. including the compds., and methods of AΒ preparation and use thereof are disclosed. The compds. are N-aryl diazaspirocyclic compds. (shown as I and II; variables defined below; e.g. III), bridged analogs of N-heteroaryl diazaspirocyclic compds., or prodrugs or metabolites of these compds. The aryl group can be a five or six-membered heterocyclic ring (heteroaryl). The compds. and compns. can be used to treat and/or prevent a wide variety of conditions or disorders, particularly those disorders characterized by dysfunction of nicotinic cholinergic neurotransmission, including disorders involving neuromodulation of neurotransmitter release, such as dopamine release. CNS disorders, which were characterized by an alteration in normal neurotransmitter release, are another example of disorders that can be treated and/or prevented. The compds. and compns. can also be used to alleviate pain. The compds. can alter the number of nicotinic cholinergic receptors of the brain of the patient, exhibit neuroprotective effects and when employed in effective amts., not result in appreciable adverse side effects (e.g. side effects such as significant increases in blood pressure and heart rate, significant neg. effects upon the gastrointestinal tract, and significant effects upon skeletal muscle). For the $\alpha 4\beta 2$ subtype, the Ki value for each of the examples of I was <1 μM , indicating that ${ t I}$ bind tightly to the receptor. Although the methods of preparation are not claimed, 13 example prepns. are included. For example, III was prepared in 5 steps (76, 93, 96, 66 and 88 % yields, resp.) starting from Et (S)-N-benzylpyrrolidine-2-carboxylate and nitroethylene and involving intermediates Et 2-(2-nitroethyl)-1-benzylpyrrolidine-2carboxylate, 6-benzyl-2,6-diazaspiro[4.4]nonan-1-one, 1-benzyl-1,7diazaspiro[4.4] nonane and 1-benzyl-7-(3-pyridyl)-1,7diazaspiro[4.4]nonane. For I: Q1 is (CZ2)u; QII is (CZ2)v; QIII is (CZ2)w; and QIV is (CZ2)x; u, v, w and x are individually 0-4, preferably 0-3; R is H, lower alkyl, acyl, alkoxycarbonyl or aryloxycarbonyl; Z is H and (un) substituted alkyl, cycloalkyl, heterocyclyl, aryl, alkylaryl, arylalkyl; Cy is a six membered ring linked via C to the N of the rest of I and each of the remaining ring atoms = N, N bonded to O or C bonded to a substituent species, wherein ≤ 3 are N or N bonded to O, or Cy is a five 5-membered heteroarom. ring linked via C to the N of the rest of I; addnl. details are given in the claims. For II: QV = (CZ2)y; QVI = (CZ2)z; y and z = 0-4; the bridged diazaspirocyclic ring contains 8-13 members; the rest of the variables are defined similarly to those for I. 646055-79-0P, Ethyl 1-azabicyclo[2.2.1]heptane-2-carboxylate 646055-80-3P, Ethyl 1-aza-2-(2-nitroethyl)bicyclo[2.2.1]heptane-2-

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of N-aryl diazaspirocyclic compds. as nicotinic cholinergic receptor modulators for treating nervous system and other disorders) 646055-79-0 CAPLUS

1-Azabicyclo[2.2.1]heptane-2-carboxylic acid, ethyl ester (9CI) (CA INDEX NAME)

RN 646055-80-3 CAPLUS

CN 1-Azabicyclo[2.2.1]heptane-2-carboxylic acid, 2-(2-nitroethyl)-, ethyl ester (9CI) (CA INDEX NAME)

L7 ANSWER 2 OF 4 CAPLUS COPYRIGHT 2004 ACS on STN

AN 1991:6240 CAPLUS

DN 114:6240

TI Synthesis of acyclic and heterocyclic derivatives of 2-carboxyquinuclidine. IV

AU Bulacinski, Andrzej Benedykt

CS Dep. Technol. Pharm. Prod., Sch. Med., Warsaw, 02-097, Pol.

SO Acta Poloniae Pharmaceutica (1989), 46(5-6), 429-34

CODEN: APPHAX; ISSN: 0001-6837

DT Journal

LA Polish

GI

CONH(CH₂)_n
$$\mathbb{R}^{1}$$
, \mathbb{R}^{2}

Quinuclidine derivs. I (n = 1, R1 = H, R2 = 2- and 4-MeO and 2-, 3-, and 4-C1; n = 2, R1 = 3-MeO, R2 = 4-MeO; R1 = H, R2 = 4-C1) were prepared in 47-72% yields by treating 2-chlorocarbonylquinuclidine.HCl with the appropriately substituted phenethylamine and benzylamine, resp., in C6H6 in presence of Et3N. Phenylalkylaminomethylquinuclidines II (n = 1, R1 = H, R2 = 2- and 4-MeO; n = 2, R1 = 3-MeO, R2 = 4-MeO) were obtained by reduction of the corresponding I with LiAlH4 in Et2O-THF.

IT 130877-60-0P 130877-61-1P 130877-65-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reduction of)

RN 130877-60-0 CAPLUS

RN 130877-61-1 CAPLUS

 $1-Azabicyclo[2.2.1] heptane-2-carboxamide, \ N-[(4-methoxyphenyl)methyl]-1-Azabicyclo[2.2.1] heptane-2-carboxamide, \ N-[(4-methoxyphenyl)methyl]-$ CN (9CI) (CA INDEX NAME)

130877-65-5 CAPLUS RN

1-Azabicyclo[2.2.1]heptane-2-carboxamide, N-[2-(3,4-dimethoxyphenyl)ethyl]-CN (9CI) (CA INDEX NAME)

130877-62-2P 130877-63-3P 130877-64-4P

130877-66-6P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

130877-62-2 CAPLUS

1-Azabicyclo[2.2.1]heptane-2-carboxamide, N-[(2-chlorophenyl)methyl]-CN (9CI) (CA INDEX NAME)

RN 130877-63-3 CAPLUS

1-Azabicyclo[2.2.1]heptane-2-carboxamide, N-[(3-chlorophenyl)methyl]-CN (9CI) (CA INDEX NAME)

130877-64-4 CAPLUS RN

1-Azabicyclo[2.2.1]heptane-2-carboxamide, N-[(4-chlorophenyl)methyl]-CN (9CI) (CA INDEX NAME)

RN 130877-66-6 CAPLUS

L7 ANSWER 3 OF 4 CAPLUS COPYRIGHT 2004 ACS on STN

AN 1984:22615 CAPLUS

DN 100:22615

TI Double cycloaddition reaction of imidazolium methylides. Intermolecular 1,3-dipolar and intramolecular Diels-Alder cycloaddition reactions

AU Tsuge, Otohiko; Kanemasa, Shuji; Takenaka, Shigeori

CS Interdiscip. Grad. Sch. Eng. Sci., Kyushu Univ., Kasuga, 816, Japan

SO Bulletin of the Chemical Society of Japan (1983), 56(7), 2073-6 CODEN: BCSJA8; ISSN: 0009-2673

DT Journal

LA English

GΙ

AB Imidazolium methylides such as imidazolium dicyanomethylide and bis(ethoxycarbonyl)methylide react with the methylenecyclopropenes with unsatd. substituents at the 4-position in the fashion of double cycloaddn. reaction, leading to the novel cage compds., e.g. I, which involves an intermol. 1,3-dipolar cycloaddn. reaction and an intramol. Diels-Alder reaction.

IT 87446-61-5P

RN 87446-61-5 CAPLUS

CN 2,4,4a-Metheno-4aH-7-oxa-1,2a-diazacyclopent[cd]indene-3,3(4H) dicarboxylic acid, 5-cyano-1,2,7a,7b-tetrahydro-1-methyl-4,6,8-triphenyl-,
 diethyl ester (9CI) (CA INDEX NAME)

LA

·=>

English

- *** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

=> d scan

L3 14 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):13

L3 14 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

IN 1-Azabicyclo[2.2.1]heptane-2-carboxamide, N-[(4-methoxyphenyl)methyl]-(9CI)

MF C15 H20 N2 O2

$$\bigcap_{C-NH-CH_2}^{O}$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L3 14 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

MF C14 H17 C1 N2 O

$$\begin{array}{c} \overset{\text{O}}{\underset{\text{C-NH-CH}_2}{\parallel}} \\ & \overset{\text{O}}{\underset{\text{C1}}{\parallel}} \end{array}$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L3 14 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

IN 1-Azabicyclo[2.2.1]heptane-2-carboxylic acid, ethyl ester (9CI)

MF C9 H15 N O2

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L3 14 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

IN 7H-3,6,6a-Metheno-3H-benzo[f]pyrrolo[3,2,1-ij]quinoline-5-carboxylic acid, 7-cyano-5,6,1lb,1lc-tetrahydro-6,12-diphenyl-, ethyl ester, (3 α ,5 α ,6 α ,6a α ,7 β ,1lb β ,1lc β ,12R*)- (9CI)

MF C32 H26 N2 O2

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L3 14 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

IN 2,4,4a-Metheno-4aH-7-oxa-1,2a-diazacyclopent[cd]indene-3,3(4H)-dicarboxylic acid, 5-cyano-1,2,7a,7b-tetrahydro-1-methyl-4,6,8-triphenyl-,diethyl ester (9CI)

MF C35 H31 N3 O5

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L3 14 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

MF C14 H17 C1 N2 O

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L3 14 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

MF C17 H24 N2 O3

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L3 14 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

MF C11 H18 N2 O4

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L3 14 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

IN 7H-3a,4,7-Methenopyrano[2,3,4-hi]indolizine-5,5(4H)-dicarboxylic acid,
3-cyano-9a,9b-dihydro-2,4,10-triphenyl-, diethyl ester (9CI)

MF C36 H30 N2 O5

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

L3 14 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

MF C15 H20 N2 O2

14 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN 1-Azabicyclo[2.2.1]heptane-2-carboxamide, N-[(3-chlorophenyl)methyl]-IN (9CI)

C14 H17 C1 N2 O MF

$$\begin{array}{c|c} \text{N} & \text{O} \\ \text{II} & \text{C-NH-CH}_2 \\ \end{array}$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

14 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN L3

1-Azabicyclo[2.2.1]heptane-2-carboxamide, N-[2-(4-chlorophenyl)ethyl]-(9CI)

C15 H19 C1 N2 O MF

$$\bigcap_{C-NH-CH_2-CH_2}^{O}$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

14 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

7H-3,6,6a-Metheno-3H-benzo[f]pyrrolo[3,2,1-ij]quinoline-5,5(6H)dicarboxylic acid, 7-cyano-11b,11c-dihydro-6,12-diphenyl-, diethyl ester, $(3\alpha, 6\alpha, 6a\alpha, 7\beta, 11b\beta, 11c\beta, 12R^*)$ - (9CI)

C35 H30 N2 O4 MF